# Texas A&M University - Corpus Christi

# **Genomics Core Lab**

# Metabarcoding Manual

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## 5 ABSTRACT



## **6 1 INTRODUCTION**

- 7 The GCL metabarcoding process involves numerous scripts in a variety of programming languages.
- 8 A metabarcoding project is done by creating a pipeline of these scripts and setting the parameters of
- 9 each. Because the needs of individual projects differs, it is difficult to create a one-size-fits-all program.
- Therefore, the use of small, modular programs appears to be the most effective. This manual describes
- each script in the GCL pipeline, demonstrates an example pipeline, and offers guidance on setting up
- databases and running on an HPC.

## 13 1.1 Charybdis

- 14 All of the code resides in a repository called GCL\_Charybdis. The repository is located on the
- HPC at /work/hobi/GCL/GCL\_Charybdis. The name resulted from finding a critter from species
- 16 Charybdis from our first set of metabarcoding results.

### 17 2 PIPELINE PARAMETERS

- 18 You need a number of files and bits of information in order to run the pipeline. The following describes
- each parameter, and (if relevant), how to obtain them. See section 4 for actual examples of these parameters
- used in a project.

## 21 2.0.1 Project Name (-p)

- 22 The name of the project is used as a prefix to many files generated throughout the pipeline. Additionally,
- 23 the input files need to use this project name in their filenames.

## 24 2.0.2 Input Directory (-i)

- <sup>25</sup> Certain files need to be placed in this directory before executing the pipeline. Instead of forcing the user
- 26 to specify several files, they are placed in a single location. The pipeline finds files by using the project
- 27 name specified with the -p option. The required files for running the generic pipeline script is shown in
- Table 1.

## **Table 1.** Files required to exist in the input directory.

The variable project name is represented by projectname>.

forward.fastq
fastq
color name > \_reverse.fastq
fastq
Tab-delimited file to match up barcodes to samples

#### 9 2.0.3 Output Directory (-o)

Files generated by pipeline will be placed in here.

### 2.0.4 Chunks (-n)

- Number of parallel threads to spawn. Use all the cores available if running on an HPC node. But if
- running on desktop, use less than your number of available cores. For the 40-core workstation in Bird's
- computer room, I would go with 35.

### 35 2.0.5 BLAST Database (-b)

- Any valid BLAST database. GCL typically uses the the nucleotide (nt) database from NCBI.
- 37 ftp://ftp.ncbi.nlm.nih.gov/blast/db/
  - Note that if you use the -b option, do not use -v option.
- See section 5.4

#### 2.0.6 BLAST Ignore File (-d)

- A text file containing GIs of sequences in the BLAST database to ignore. It is fine to have GIs that are not
- actually in the BLAST database used. Good for filtering environmental DNA and such.

#### 43 2.0.7 VSEARCH Database (-v)

- VSEARCH is an alternative taxonomic assignment program Rognes et al. (2016). A valid VSEARCH
- database is a FASTA file. But because FASTA files are a loose standard, not all are compatible. VSEARCH
- 46 gets unhappy regarding certain formats of the sequence header. An example of a valid FASTA entry is
- as follows (data is from (Ratnasingham and Hebert, 2007). Note that the | symbol is used to separate
- metadata field in the header. The first field is a unique sequence ID. The second is a scientific name. The
- 49 third is the genetic region of the sequence. The fourth is the Barcode of Life Database (BOLD) sequence
- ID. It could be a NCBI GI or Accession instead.

Note that if you use the -b option, do not use -v option.

#### See section 5.4

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>GBMAA467X14| Pomphorhynchus\_laevis | COIX5P | KF559289 ATGTATGTTTGGTTGGTGTGAGGGGGGCTAATGGGGTTTTCTATAAGACTATTAATTCGA

The script /work/hobi/GCL/GCL\_Charybdis/bin/vsearch\_getTAXIDfromBOLDseqid.sh will get the NCBI taxonomic ID from a BOLD sequence ID.

### 7 2.0.8 Chimera Database (-c)

- 58 FASTA database used for detection on chimeric reads. Chimera detection can be done using a database or
- using the de novo method Rognes et al. (2016). Currently the pipeline is hard-coded to use the de novo
- 60 method, but still requires a chimera database that will not be used. This issue is described in section 5.3.

### 2.0.9 Target Sequence Length (-x)

- 62 The target sequence length is the length of a read expected using PCR. The forward and reverse primers
- extract a portion of DNA between them. The value of this option is based on the primers you used, so
- check their documentation or ask whoever did the PCR.

### 55 2.0.10 Charybdis Scripts Directory (-g)

- 66 Instead of setting the environment \$PATH variable or placing the scripts in a systemwide bin directory,
- 67 the directory holding all the pipeline scripts is an option. Less conventional than \$PATH-based methods,
- but very easy to deal with.

## 2.0.11 NCBI Taxonomy Database (-t)

- This database is used to get an organism's scientific name from a numeric taxonomic ID. Taxonomy
- database downloaded from ftp://ftp.ncbi.nlm.nih.gov/pub/taxonomy/.
- 72 It is the same data as https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi

## 3 PIPELINE ARCHITECTURE

- 74 Pipelines are bash scripts that just string together a group of other scripts/programs to perform the metabar-
- 75 coding process. Example scripts are included in /work/hobi/GCL/GCL\_Charybdis/pipelines.
  - The script called charybdis\_generic is expected to handle most projects with little customization.
- The section will describe all of the steps of the charybdis\_generic pipeline. Hopefully others will be able to use this information to create a pipeline tailored to other metabarcoding projects.

- Throughout the text, *attribute* refers to a piece of metadata included in a FASTA sequence header.
- 80 These are in the form of key-value pairs. Many tools in the OBITOOLS package create and modify
- attributes, so we made many of our own scripts follow this convention.

## 82 3.1 Merge Reads

- 83 Script: /work/hobi/GCL/GCL\_Charybdis/bin/mergeReads.sh
- 84 Slurm wrapper: /work/hobi/GCL/GCL\_Charybdis/bin/mergeReads.slurm
- The forward and reverse reads and merged into a single FASTA file, and basic quality filtering is done.
- The reads are matched to sample ID which are included as metadata in the FASTA sequence headers.

### 87 3.1.1 Split FASTQ files

- 88 In order to run the next sections in parallel, the forward and reverse read FASTQ files are split into equally
- sized chunks using fastq-splitter.pl.

#### 90 3.1.2 Join forward and reverse reads

- 91 After this point, the pipeline uses a single file of merged forward and reverse reads. This is done using
- 92 OBITOOL's illuminapairedend tool. We have it reject reads whose alignment score score is under a
- 93 threshold of 40.

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#### 94 3.1.3 Convert FASTQ to FASTA

- 95 Convert the merged files to FASTA. This is done because many tools operate only on FASTA and because
- 96 we don't need the sequencer quality information after this point. We can always use the unique sequence
- 97 ID to access it from the origin FASTQ, if needed.

### 3.1.4 Remove unaligned sequences

- <sup>99</sup> We use OBITOOL's *obigrep* to select only those sequences whose alignment mode is "joined". *Obigrep*
- can search for patterns of values of specified attributes.

### 3.1.5 Remove sequences whose alignment length is below threshold

- We use OBITOOL's *obigrep* to select only those sequences whose alignment length is  $\geq$  a threshold. We
- have this threshold set to 20.

#### 3.1.6 Match sequences to sample ID based on barcode

- OBITOOL's ngsfilter is used to attach an attribute called sample to each sequence. The match is allowed
- to have 2 sequence errors by setting the flag -e 2.

#### of 3.1.7 Concatenate into single FASTA

The small chunks are concatenated into a single FASTA file.

#### 109 3.2 Filter Reads

- Script: /work/hobi/GCL/GCL\_Charybdis/bin/ObiToolsPipeline.sh
- Slurm wrapper: /work/hobi/GCL/GCL\_Charybdis/bin/filterReads.slurm

This was the first script written. At the time, the size of the project was not known and this file was intended to be the entire pipeline. Thus, the script is called <code>ObiToolsPipeline.sh</code> instead of the more appropriate <code>filterReads.sh</code>.

## 3.2.1 Split into smaller files by sample ID

As before, we form a smaller number of files for parallel processing. However, the sample ID is used to split files instead of an arbitrary size. The sample ID is added to the FASTA filename to keep track.

## 3.2.2 Remove unneeded attributes

OBITOOLS added a huge number of attributes in a previous step. We get rid of the irrelevant ones with obiannotate. Note that attributes are never truly discarded since the sequence ID can always be used to query the intermediate FASTAs generated at each step of the pipeline.

## 3.2.3 Keep only unique sequences

In order to lower the filesize (often substantially), duplicated sequences are merged into a single representative sequence. The *obimerge* attribute is used to record how many sequences are represented by that sequence. This is done using obiuniq.

#### 3.2.4 Remove PCR errors

PCR errors are filtered using obiclean. After numerous experiments, we found that the following options were performing well for a set of COI sequences from Gulf of Mexico fish. Results from projects where we reused these settings have appeared to be fine.

## **Table 2.** Settings for obiclean

- -r 0.5 Controls when less abundant sequences are labeled as variants of a similar more abundant
- -d 1 Max number of differences to be considered variants
- -H Keep only a representative of a set of variants.

#### 3.2.5 Remove low read length

The amplified sequence should be a certain size. Using our COI primers, we expect the size to be 313 base pairs. However, the actual size is often a bit lower or higher. You supply your base pairs length as a pipeline argument. The allowed range is that size  $\pm$  15.

#### 3.2.6 Remove chimeras

VSEARCH is used to remove chimeras using the *de novo* method.

### 36 3.2.7 Concatenate into single FASTA

The small chunks are concatenated into a single FASTA file.

## 3.3 Collapse into OTUs

Script: /work/hobi/GCL/GCL\_Charybdis/bin/ClusterOTU.sh

140 Slurm wrapper: /work/hobi/GCL/GCL\_Charybdis/bin/ClusterOTU.slurm

### 141 3.3.1 Cluster with CROP

CROP clusters sequences into OTUs (Hao et al., 2011). CROP has a number of parameters that control the OTU assignment. Assigning these parameters is tricky, but the CROP manual has a recommended process for determining the values (TingChenLab, 2017). We initially used the recommendation, but then modified it to get better results. However, we don't currently remember how we figured out some of the values used in our process. You can check the manual and compare to ours, but note that -*b* and -*z* should only affect performance, not clustering. However, we set -*e* based on -*z* (following the manual), which does affect results. We need to investigate further.

#### **Table 3.** Settings for CROP.

*NUMSEQS* is the total number of sequences. *BP* is the target sequence length.

-r	5	Clusters of size <i>leqr</i> are considered rare
-S		Specifies that 97% similarity needed to cluster
-b	NUMSEQS/50	Number of blocks
-Z	(150000/BP) - 0.1 * (150000/BP)	Max number of sequence a block can hold
-е	[-z]*10	Number of MCMC iterations

#### 9 3.3.2 Correct the OTU size

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150 CROP records what sequences end up in an OTU, as well as the total number of sequences represented 151 by an OTU. However, CROP has no knowledge that objclean and objmerge already collapsed 152 duplicate/similar sequences into. We call CROP\_size\_fix.sh to get the actual OTU counts.

## 3.4 Taxonomic Assignment

Taxonomic assignment can be done using either BLAST or VSEARCH. (SAP coming soon). The exact scripts and slurm wrappers depend on method selected.

## 3.4.1 Run taxonomic assignment program in parallel

Split sequences into chunks and BLAST in parallel. Uses the BLAST ignore list (provided in parameters) to skip assignment to unwanted sequences. For example, we typically try to avoid assignment to environmental sequences.

#### 3.4.2 Concatenate into single FASTA

The small chunks are concatenated into a single FASTA file.

#### 3.4.3 Convert to Charon format

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163 Charon is the name of our own CSV format for assigned sequences. The purpose is that we can write
164 conversion scripts to have assignments from various taxonomic assignment programs into a standard. The
165 Charon file has columns specified by Table 4.

**Table 4.** Columns for Charon CSV file

Query Sequence ID Assignment Sequence ID Scientific Name NCBI GI Number of sequences in OTU

# 3.5 OTUs VS Samples

Formerly: Critters VS Tubes.

Will explain soon, but basically just a script that makes a column for each sample and a row for each OTU. The cells have a count of the number of sequences of that OTU in that sample. Also has columns of the scientific names for various phylogenetic levels. Also has extra information columns such as the sequence, BLAST scores, etc. No limit on added extra columns to this CSV file.

### 4 EXAMPLE RUN

## 4.1 Prepare Data

## 4.2 Run Pipeline Script

The following bash script executes charybdis\_generic.sh, an implementation of the GCL pipeline.
The code segment shown is a single line of bash, using \ for clarity to break it into one line per option.

```
bash /work/hobi/GCL/GCL_charybdis/pipelines/charybdis_generic.sh \
178
179
       -p Simons-H1_ATTACTCG-TAATCTTA_L001 \
       -i /work/hobi/GCL/20180330 - Simons/Simons-H1/in \
180
       -o /work/hobi/GCL/20180330-Simons/Simons-H1/out
181
       -n 20
182
       -b /work/hobi/GCL/db/NCBI_BLAST_DBs/nt \
       -d /work/hobi/GCL/db/GI_lists/environmental.NCBI_nucl__SORTED.gi \
184
       -c /work/hobi/GCL/db/BOLD_FULL.fasta \
          313 \
186
          /work/hobi/GCL/GCL_charybdis/bin \
187
       -t /work/hobi/GCL/db/TAXO
188
```

Each option will be explained, and the first 10 lines of each file will be displayed. This way, you can compare the format of your input files to those used for the example. Do not trust the spacing seen here. Tabs may become spaces, etc. See the official file format specifications.

-p Name of project.

We used Simons-H1\_ATTACTCG-TAATCTTA\_L001 to match the names of the input FASTQ files. The important thing is that each project should have a unique name for keeping projects separate/organized.

-i Input directory.

This is where the input files should be placed. A slight abuse of terminology, since projectname>.samples.txt
is generated by the pipeline, by reading sample IDs from projectname>.barcodes.txt
Note that the FASTQ sequences were shortened to fit on the page. A sequence of five periods (....) represents a removed segment.

```
{\tt ACTTGATAGACCTCGGGGTGGCCGAAGAACCAGAATAGGTGTTGGTAAAGAATTGGGTCTCC}.....
208
209
        210
        @MISEQ01:242:000000000 - BLT5R:1:1101:16137:1376 1:N:0:ATTACTCGTAATCTTA
211
        ATCACGGGAACTGGATGAACAGTTTATCCTCCCCTTGCCGGCAACCTGGCCCACGCAGGGGC\ldots
212
213
       214
        @MISEQ01:242:000000000 - BLT5R:1:1101:15575:1386 1:N:0:ATTACTCGTAATCTTA
215
        \tt GTCCGCGGTACTGGATGAACAGTATACCCCCCTTTAGCAGCAGCCATTGCACATGCAGGTGC\ldots ...
216
217
    [ekrell@hpcm Simons-H1] head in/Simons-H1_ATTACTCG-TAATCTTA_L001_forward.fastq
218
        @MISEQ01:242:000000000 - BLT5R:1:1101:15746:1350 1:N:0:ATTACTCGTAATCTTA
219
        ACTTGATAGACCTCGGGGTGGCCGAAGAACCAGAATAGGTGTTGGTAAAGAATTGGGTCTCCCCCA\dots
220
221
       222
        @MISEO01:242:000000000 - BLT5R:1:1101:16137:1376 1:N:0:ATTACTCGTAATCTTA
223
        ATCACGGGAACTGGATGAACAGTTTATCCTCCCCTTGCCGGCAACCTGGCCCACGCAGGGGCATCA\dots \\
224
       226
        @MISEQ01:242:000000000 - BLT5R:1:1101:15575:1386 1:N:0:ATTACTCGTAATCTTA
       \tt GTCCGCGGTACTGGATGAACAGTATACCCCCCTTTAGCAGCAGCCATTGCACATGCAGGTGCATCT\dots...
228
229
   [ekrell@hpcm Simons-H1]$ head in/Simons-H1_ATTACTCG-TAATCTTA_L001.barcodes.txt -n 5
230
                     sample
        #exp
                                      forward_primer reverse_primer
                              tags
231
                             ACAGTG: CCGTCC
        Simons-H1
                     1901 - B
                                              GGWACWGGWTGA . . . . .
                                                                         TANACYTCNGGRTGN
232
        Simons-H1
                     1685-C
                             ACAGTG: GTTTCG
                                              GGWACWGGWIGA....
                                                                         TANACYTCNGGRTGN . . . .
233
        Simons-H1
                     1759-B
                             ACTGAT: ATGTCA
                                              GGWACWGGWTGA . . . . .
                                                                         TANACYTCNGGRTGN . . . . .
234
        Simons-H1
                     1774-X ACTGAT: GCCAAT
                                               GGWACWGGWTGA . . . . .
                                                                         TANACYTCNGGRTGN . . . . .
235
      -o Output directory.
236
   Files generated by the pipeline, including numerous intermediate file, are placed here.
237
      -n Number of parallel instances.
238
   The nodes on TAMUCC's HPC cave 20 cores, so 20 were used.
239
      -b BLAST Database.
240
   Specify path to BLAST database. We are using the nucleotide (nt) database,
241
   which is downloaded from ftp://ftp.ncbi.nlm.nih.gov/blast/db/.
242
      -d List of GIs for BLAST to ignore This is a list of GIs, where each GI specifies a sequence in the BLAST
243
   database to ignore. This file is mandatory, but may be blank (See Section 5.2).
244
   We attempt to remove environmental DNA, so our list has GIs of environmental sequences.
    [ekrell@hpcm Simons-H1]$ head -n 5 ...../environmental.NCBI_nucl__SORTED.gi
246
      1000014437
247
      1000014439
248
      1000014441
249
      1000014443
250
      1000014445
251
      -c Chimera Database. We use a FASTA file generated from the BOLD database. But, we ended up doing de novo
252
   chimera detection. Currently this file is required and just not being used (See section 5.3).
253
      -x Target sequence length (basepairs). This number is based on the primers used during PCR.
254
      -g Directory of charybdis scripts.
255
    ls /work/hobi/GCL/GCL_charybdis/bin
256
                                         fastq-splitter.pl
257
      AddBlast.slurm
                                         filterReads.slurm
258
      AddVsearch.slurm
                                        mergeReads.sh
259
      AssignToMarkers.sh
                                        mergeReads.slurm
260
      AssignToMarkers.slurm
                                         ObiToolsPipeline.sh
261
      blast_10custom_to_charon_OTU.R
                                        ObiToolsPipeline.slurm
262
                                        ObiToolsPipeline_stats.sh
      BlastMeta.sh
263
      BlastMeta.slurm
                                        OTU_CVT_addBlast.R
264
      ClusterOTU.sh
                                        OTUvsTube.slurm
265
```

```
ClusterOTU.slurm
                                           split_by_marker.R
266
      CombineAndCROP.sh
                                           vsearch_blast6custom_to_charon -BLASTDB_OTU.R
267
      critters V Stubes _ OTU . R
                                           vsearch_blast6custom_to_charon_OTU.R
268
                                           vsearch\_getTAXIDfromBOLDseqid.sh
      CROP_size_fix.sh
      CROP. slurm
                                           Vsearch.sh
                                           Vsearch.slurm
      determine_marker.R
      determine_marker.sh
272
       -t Directory with NCBI taxonomy database Taxonomy database downloaded from ftp://ftp.ncbi.nlm.
273
   nih.gov/pub/taxonomy/.
274

    g Directory of charybdis scripts.

275
    [ekrell@hpcm_pipelines]$ ls /work/hobi/GCL/db/TAXO
276
             Accesion2Taxid delnodes.dmp
                                                              merged.dmp
                                                                            nodes.dmp
                                                                                          taxdump.tar.
277
                                               gc.prt
             citations.dmp
                               division.dmp
                                               gencode.dmp
                                                              names.dmp
                                                                            readme.txt
278
```

# 5 AREAS FOR IMPROVEMENT

## 5.1 Support NCBI Accession IDs

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NCBI is phasing out GIs (sequence IDs), which this pipeline uses at various steps (NCBI, 2016). Accession IDs are the current standard.

#### 283 5.2 Pipeline should not require BLAST ignore list

<sup>284</sup> Currently this is required. If you don't need it, you can supply a blank file. Not very elegant. Would be simple to make it optional.

#### 5.3 Chimera database should not be mandatory

The generic pipeline uses *de novo* chimera detection, but still requires the chimera database. A better solution would be to default to *de novo*, but use the database if supplied.

#### 5.4 Taxonomic assignment methods need not be mutual exclusive

Currently, you have to specify either a BLAST or VSEARCH database. Will receive a warning if attempting to supply both options. But, there is no reason to keep the user from specifying both. The pipeline could simply branch at that point, do assignment with both, and perform remaining pipeline steps with both results. Work would need to be done to organize filenames to differentiate. The current problem is that all assignment methods generate files with the same name. The assignment method itself would need to be part of the filename.

# 6 STATISTICAL ASSIGNMENT PACKAGE (SAP)

### We have SAP working, but not meshed into the pipeline yet. So it has its own section, at least for now.

SAP (Munch et al., 2008) differs from typical assignment software (BLAST, VSEARCH) in that it is based on computing a posterior probability that the query sequence is a member of a particular clade. Two major phases are involved in an SAP taxonomic assignment. First, BLAST is executed to find a set of similar sequences. Since similar sequences should indicate similar biological properties, the set of accepted BLAST sequences are called the set of homologues with the query. To strengthen the assumption that the sequences are homologous, there is a limit on how distant the furthest apart homologues can be. Several parameters are used to control BLAST's sequence search and also to control how SAP chooses which of those sequences to include in the homologue set. It can and does happen that a single species has several sequences in the database. However, a particular taxonomic group appears only once in the phylogenetic tree. Therefore, SAP focuses on diversity and only uses the best match for a given species. While this is necessary to construct meaningful trees, it also means that the species which only appears once in a set of BLAST results is given equal weight to a predominate species result. The set of homologues and the query sequence are used to sample a large number of trees using Markov Chain Monte Carlo. The posterior probability for any taxonomic group is based on the proportion of trees in which the query sequence was in that group. A simple case that shows how this can differ from BLAST occurs when several species from the same genus are present in the homologue set. Where BLAST would simply list the hits in order of sequence ID, SAP determines that it cannot reliably be determined what species the query belongs to. A top hit may differ little from the hits just below it, and it is essentially arbitrary to assume that the top hit is the exact species when it scored closely against other species as well. The SAP result would give a low posterior probability to any particular sequence, but the genus would have a very high probability.

The above text taken from GCL's work-in-progress manuscript "An Evaluation of Software for Taxonomic Assignment with DNA Metabarcoding"

- 6.1 Fixing SAP
- 319 6.2 Running SAP
- 320 6.3 Evaluating SAP
- 321 6.3.1 Categorize SAP results
- 6.3.2 Generate SAP report
- 323 6.4 SAP outstanding issues
- 324 6.5 Complete SAP example

## 325 REFERENCES

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